Activating transcription factor 3 inhibits endometrial carcinoma aggressiveness via JunB suppression

FANGYUAN WANG1, JINGJING LI2, HAIXIA WANG3, FAN ZHANG4 and JIN GAO2,4

1Shanghai Institute of Rheumatology, Renji Hospital, School of Medicine, Shanghai Jiao Tong University, Shanghai 200127; 2School of Pharmacy; 3School of Agriculture and Biology, Shanghai Jiao Tong University, Shanghai 200240, P.R. China; 4Department of Pharmaceutical Sciences, College of Pharmacy and Pharmaceutical Sciences, Washington State University, Spokane, WA 99202, USA

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Abstract. The function of activating transcription factor 3 (ATF3) in cancer is context-dependent and its role in endometrial carcinoma (EC) is yet to be elucidated. In the present study, ATF3 was indicated to be downregulated, while one of the ATF3-interacting proteins, JunB, was upregulated in ECs according to western blot analysis. After overexpression in ECs, ATF3 inhibited the proliferation and invasion of EC cells and enhanced apoptosis, as well as suppressed the expression of JunB. The properties of EC cells, including the expression of matrix metalloproteinases, tissue inhibitors of metalloproteinases, the cell cycle and apoptosis were all altered by overexpression of ATF3. Furthermore, luciferase activity assay, chromatin precipitation and DNA affinity assay results indicated that ATF3 exerted the aforementioned functions via JunB binding and activator protein-1 signaling. However, the interaction between ATF3 and JunB did not occur in EC cells under basal conditions, but in ATF3-overexpressing ECs, which was capable of mitigating EC proliferation, invasion and metastasis. Collectively, the present results suggested that the ATF3/JunB interaction may serve as a potential therapeutic target for ECs.

Introduction

Endometrial carcinoma (EC) is one of the most common types of gynecologic malignancy, with an estimated 61,880 diagnosed cases, accounting for 6.94% of all new female cancer cases and 12,160 mortalities, leading to 4.26% of all mortalities in females in 2019 according to the American Cancer Society (1). With the development of medical technologies, the majority of ECs are diagnosed at the early stage (stage I-II), according to staging standards of the International Federation of Gynecology and Obstetrics, and hysterectomy is the recommended treatment (2). However, 30% of EC cases are not diagnosed until metastasis is identified (stage III-IV), which is associated with low survival and worse prognosis (3,4). Furthermore, particularly for patients with EC who are nulliparous, hysterectomy is not a primary treatment option (5). Currently, the global incidence of EC is on the rise. In 2013, there were an estimated 49,560 cases in the USA, while by 2018, the number of new cases had risen to 63,230 individuals (6); this increase may be attributed to an increased prevalence of patients who are overweight, obese and physically inactive (7,8). Therefore, therapeutic strategies focusing on inhibiting the proliferation, migration and invasion of EC are critical to the treatment and prognosis of patients with EC. However, the molecular and cellular mechanisms responsible for the carcinogenesis, progression and survival prognosis of ECs have remained elusive.

EC is classified into two subtypes: Type I, accounting for 85% of total cases, 20-50% of which have acquired p53-mutant status; and type II, accounting for 15% of total cases, >90% of which have p53-mutant status (9-11). Previous studies have reported that activating transcription factor 3 (ATF3) binds to the C terminus of wild-type (WT) p53, preventing p53 from ubiquitin-mediated degradation in response to DNA damage (12). Moreover, ATF3 is able to bind to mutant p53 to inhibit the migration and invasion of p53-mutated cancer cells, suppressing the oncogenic function of mutant p53 proteins (13). Therefore, ATF3 may be a potential therapeutic target for carcinomas closely linked to p53 mutations, including ECs. To the best of our knowledge, the present study was the first to investigate the role of ATF3 in EC. Its implication in cell proliferation, invasion, signaling pathway markers, matrix metalloproteinases (MMPs), tissue inhibitors of metalloproteinases (TIMPs), protein interaction and DNA binding were assessed to identify the regulatory function of ATF3 in ECs and the potential underlying mechanisms.
Materials and methods

Ethics statement. The protocol (approval no. 0101N16032) of the present study was approved by the Human Investigation Ethical Committee of Shanghai First People's Hospital Affiliated Shanghai Jiao Tong University (Shanghai, China) and Renji Hospital Affiliated to Shanghai Jiao Tong University School of Medicine (Shanghai, China). The sampling methods in patients were performed in accordance with the Operational Guidelines for Ethics Committees That Review Biomedical Research (https://www.who.int/trd/publications/documents/ethics.pdf). All samples were obtained from patients who had provided written informed consent for the use of their tissues for the purposes of research after the operation.

The animal experiment was in accordance with the recommendations in the Guidelines for the Care and Use of Laboratory Animals of China (http://www.nsfc.gov.cn/nsfc/cen/cen/pfzl/pufanew/20110801_02.pdf). The protocol (approval no. 2013020346) was approved by the Committee on the Ethics of Animal Experiments of the School of Pharmacy of Shanghai Jiao Tong University (Shanghai, China). All efforts were made to minimize animal suffering.

Cell culture, plasmids and transfections. The human EC cell lines HEC-1B and AN3CA were cultured in DMEM/nutrient mixture F-12 (DMEM/F12; Hyclone; Cytiva) supplemented with 10% heat-inactivated fetal bovine serum (FBS, Gibco; Thermo Fisher Scientific, Inc.) and 1% penicillin/streptomycin (Gibco; Thermo Fisher Scientific, Inc.). These cell lines were purchased from the Chinese Academy of Sciences Committee Type Culture Collection and were routinely maintained in the Fangyuan Wang's laboratory. Cell cultures were maintained at 37°C in a humidified incubator with 5% CO₂. The plasmid pShuttle Vector containing the open reading frame (ORF) of ATF3 was obtained from GeneCopoeia, Inc.

For transfection, the ATF3 ORF was inserted to the multiple cloning sites of pCMV-script (Agilent Technologies, Inc.) and transfected using Lipofectamine® 3000 (Invitrogen; Thermo Fisher Scientific, Inc.). To stably express ATF3 in HEC-1B and AN3CA cells, the EC cells were transfected with pCMV-script-ATF3 or the negative control (NC) vector pCMV-script (5 μg) and then selected with G418 (800 μg/ml; Sigma-Aldrich; Merck KGaA) after 6 weeks to obtain the stable transfection cell lines. The cell lines with ATF3 overexpression were referred to as ‘HEC-1B-ATF3’ and ‘AN3CA-ATF3’, respectively, with their control cells referred to as ‘HEC-1B-NC’ and ‘AN3CA-NC’, respectively.

Tissue collection. EC samples (n=50; age, 29-67 years) and the healthy samples adjacent to tumor tissues (n=6; age, 35-62 years) were obtained from patients who received surgical therapy at the Department of Obstetrics and Gynecology, Shanghai First People's Hospital Affiliated to Shanghai Jiao Tong University (Shanghai, China), between February 2010 and December 2013. The healthy tissues were from the same group of patients for the EC tissues. In brief, a vaginal speculum was used to expose the cervix after general or local anesthesia, followed by disinfection of the cervix using 0.5% povodine iodine (Qingdao Hainuo Biology Engineering Co., Ltd.) and paralysis of the cervical canal via a multiple-point injection of 1% lidocaine 20-100 mg (Tianjin Zhongxin Pharmaceutical Group Co., Ltd.). The cervix was then dilated, and the uterine cavity was penetrated using a long and thin metal rod with a curette at the end to perform curettage of the endometrium. Tumor staging and histological grading were performed according to the International Federation for Obstetrics and Gynecology criteria (2009) (14). Clinical and pathological data of the cohort are presented in Table I.

Primary cell culture. Fresh clinical samples of the tumor (1 cm) were minced into 2-mm fragments and then transferred to a solution of trypsin 0.25% in medium (containing 0.1 mM EDTA), followed by incubation at 37°C for 20 min. Isolated EC cells were recovered in 0.5% trypsin inhibitor (Merck KGaA) and 1.0% albumin (Sigma-Aldrich; Merck KGaA) in DMEM/F12 medium and then isolated by centrifugation at 400 x g for 10 min at 4°C. The cells obtained were manually counted using a hemocytometer under a light microscope (magnification, x40; Eclipse Ts2; Nikon Corporation) and transferred to 75-cm² culture flasks and incubated in medium supplemented with 10% FBS in an incubator at 37°C with a humidified atmosphere containing 5% CO₂.

Immunohistochemical staining. EC tissues, adjacent healthy tissues or xenografts were fixed using 10% neutral paraformaldehyde at room temperature for 24 h and then sections with a thickness of 5 μm were prepared. After deparaffinization and dehydration, sections were boiled in 10 mM tris(hydroxymethyl) aminomethane-Ethylendiaminetetraacetic acid (Tris-EDTA) buffer to retrieve antigens and then blocked in 5% normal goat serum (Sigma-Aldrich; Merck KGaA) at room temperature for 2 h. After incubation with primary antibodies overnight at 4°C, sections and secondary antibodies at room temperature for 1 h, the sections were stained using an ABC Elite kit (cat. no. PK6200; Vector Laboratories, Inc.) and a diaminobenzidine kit (cat. no. SK4100; Vector Laboratories, Inc.) according to the manufacturer's protocol. The images of sections were captured under a light microscope (magnification, x40; Eclipse Ts2; Nikon Corporation). Primary antibodies used in the present study were rabbit polyclonal anti-ATF3 antibody (cat. no. ab87213; 1:200; Abcam), mouse monoclonal anti-JunB antibody (cat. no. sc8051; 1:200; Santa Cruz Biotechnology, Inc.) anti-proliferating cell nuclear antigen antibody (PCNA; cat. no. ab29; 1:100; Abcam) and rabbit polyclonal anti-Ki67 antibody (cat. no. ab15580; 1:100; Abcam). For the biotinylated horseradish peroxidase anti-rabbit or anti-mouse secondary antibodies were included in the ABC Elite kit; an incubation with biotinylated horseradish peroxidase anti-rabbit or anti-mouse secondary antibodies were included in the ABC Elite kit, and a dilution of 1:3,000 was used.

Reverse transcription-quantitative PCR (RT-qPCR). Total RNA was extracted from HEC-1B-NC, HEC-1B-ATF3, AN3CA-NC and AN3CA-ATF3 cells with TRIzol® reagent (Invitrogen; Thermo Fisher Scientific, Inc.) and reverse transcribed at 42°C for 30 min to cDNA using the Prime Script RT reagent kit (Takara Bio, Inc.). RT-qPCR was performed using SYBR Premix Ex Taq (Takara Bio, Inc.) and analyzed with an ABI Prism 7000 Sequence Detection System (Applied Biosystems; Thermo Fisher Scientific, Inc.). The thermocycling conditions were as follows: Initial denaturation at 95°C for 10 min, followed by 42 cycles at 95°C for 15 sec, 60°C...
Membranes were blocked using 5% non-fat milk powder and transferred to a polyvinylidene difluoride membrane. A total of 50 µg protein was separated by 12% SDS-PAGE and the protein concentrations were determined using the bicinchoninic acid (BCA) method (Pierce; Thermo Fisher Scientific, Inc.).

The primary EC cells, HEC-1-NC, HEC-1B-ATF3, AN3CA-NC or AN3CA-ATF3 cells were lysed and the protein concentrations were determined using BCA assay. The cell lysates were subject to WB analysis as aforementioned using the primary antibodies anti-poly(ADP-ribose) polymerase (PARP; 1:500; cat. no. 614302; Biolegend, Inc.), anti-Bcl2 (1:1,000; cat. no. 658702; Biolegend, Inc.), anti-Bcl-x (1:500; cat. no. 633902; Biolegend, Inc.), anti-Bax (1:200; cat. no. 633602; Biolegend, Inc.), anti-p-p38 (1:500; cat. no. 622402; Biolegend, Inc.) and anti-phosphorylated (p)-p38 antibodies (1:200; cat. no. 690202; Biolegend, Inc.).

For the apoptosis marker detection, HEC-1B-NC, HEC-1B-ATF3, AN3CA-NC or AN3CA-ATF3 cells were lysed and the protein concentrations were determined using BCA assay. The cell lysates were subject to WB analysis as aforementioned using the primary antibodies anti-poly(ADP-ribose) polymerase (PARP; 1:500; cat. no. 614302; Biolegend, Inc.), anti-Bcl2 (1:1,000; cat. no. 658702; Biolegend, Inc.), anti-Bcl-x (1:500; cat. no. 633902; Biolegend, Inc.), anti-Bax (1:200; cat. no. 633602; Biolegend, Inc.), anti-p-p38 (1:500; cat. no. 622402; Biolegend, Inc.) and anti-phosphorylated (p)-p38 antibodies (1:200; cat. no. 690202; Biolegend, Inc.).

Western blot (WB) analysis. The primary EC cells, HEC-1B and AN3CA cells were lysed using Extraction and Quantification ProteoJET Mammalian Cell Lysis reagent (Fermentas; Thermo Fisher Scientific, Inc.) with protease inhibitor cocktail (Roche Diagnostics). Total protein concentrations were determined using the bicinchoninic acid (BCA) method (Pierce; Thermo Fisher Scientific, Inc.). A total of 50 µg protein was separated by 12% SDS-PAGE and transferred to a polyvinylidene difluoride membrane. Membranes were blocked using 5% non-fat milk powder for 1 h at room temperature, incubated with primary antibodies overnight at 4˚C and then washed three times in Tris-buffered saline (pH 7.4). Subsequently, membranes were incubated for 2 h with secondary antibodies at room temperature. The primary antibodies used were as follows: Rabbit anti-ATF3 (1:200; cat. no. ab216569; Abcam), mouse anti-JunB (1:200; cat. no. sc-8051; Santa Cruz Biotechnology, Inc.) and rabbit anti-GAPDH (1:20,000; cat. no. ab199553; Abcam). Secondary horseradish peroxidase-conjugated goat anti-rabbit (1:1,000; cat. no. sc-2004; Santa Cruz Biotechnology, Inc.) and goat anti-mouse antibodies (1:4,000; cat. no. SA00001-1; ProteinTech Group, Inc.) were visualized with the Super Signal West Pico chemiluminescent substrate (Thermo Fisher Scientific, Inc.) using a gel imaging system with a preconfigured Image Lab software 5.2.1 (ChemiDoc XRS+; Bio-Rad Laboratories, Inc.).

For the apoptosis marker detection, HEC-1B-NC, HEC-1B-ATF3, AN3CA-NC or AN3CA-ATF3 cells were lysed and the protein concentrations were determined using BCA assay. The cell lysates were subject to WB analysis as aforementioned using the primary antibodies anti-poly(ADP-ribose) polymerase (PARP; 1:500; cat. no. 614302; Biolegend, Inc.), anti-Bcl2 (1:1,000; cat. no. 658702; Biolegend, Inc.), anti-Bcl-x (1:500; cat. no. 633902; Biolegend, Inc.), anti-Bax (1:200; cat. no. 633602; Biolegend, Inc.), anti-p-p38 (1:500; cat. no. 622402; Biolegend, Inc.) and anti-phosphorylated (p)-p38 antibodies (1:200; cat. no. 690202; Biolegend, Inc.).

Yeast two-hybrid (Y2H) assay. The Y2H assay was performed using a Matchmaker Gal4-based two-hybrid assay kit (Clontech Laboratories, Inc.) according to the manufacturer's instructions. A bait protein (ATF3) was expressed in the yeast strain Y2HGold as a fusion to the Gal4 DNA-binding domain (DNA-BD) using the pGBKTT7 vector with the ORF of ATF3 inserted, while the prey proteins from the the high-complexity EC library, which are provided in yeast strain Y187, were expressed as fusions to the Gal4 activation domain (AD) using the pGADT7 vector, according to manufacturer's protocols. When cultures of the two transformed strains were mixed together for 20-24 h at 30˚C with slow shaking at 1-5 x g, they mated to create diploids. Thus, the DNA-BD and AD were guided into proximity to activate the transcription of four independent reporter genes (AUR1-C, ADE2, HIS3 and MEL1, coding for enzyme inositol phosphoryl ceramide synthase, histidine, adenine and α-galactosidase, respectively). Via streaking with toothpicks onto high-stringency selective synthetic dropout media supplemented with 20 µg/ml X-α-Gal, screening was performed based on whether bait (ATF3) and prey (library) fusion proteins interacted with each other. The interactions between the baits and preys were determined 24 h after culturing at 30˚C. The positive control mating between Y2HGold [pGBKTT7-Lam] and Y187 [pGADT7-T], and the NC mating between Y2HGold [pGBKTT7-Lam] and Y187 [pGADT7-T] were performed simultaneously. Autoactivation of reporter genes in Y2HGold by the bait (ATF3) was confirmed via streaking the plasmid-harbored yeast cells on the selective synthetic dropout media prior to the two-hybrid screen, in the absence of a prey protein.

Polyhistidine (His) tag-fused ATF3 (His-ATF3) expression and purification. The ORF of ATF3 was subcloned into the

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<th>Table I. Clinicopathological summary of the samples from patients with endometrial carcinoma.</th>
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<td><strong>Clinical pathological variables</strong></td>
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<td><strong>Age (years)</strong></td>
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<td>FIGO, International Federation of Gynecology and Obstetrics (14).</td>
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pET32a vector (Novagen; Merck KGaA) and introduced into Escherichia coli BL21 (Invitrogen; Thermo Fisher Scientific, Inc.) following the manufacturer's protocol. His-ATF3 was induced in the host bacteria using 0.2 µM isopropyl-β-D-1-thiogalactopyranoside (IPTG; Sigma-Aldrich; Merck KGaA) at 37°C for 4 h and the soluble protein of His-ATF3 was purified using a His-Tag purification column (Sigma-Aldrich; Merck KGaA) according to the manufacturer's instructions.

**BIAcore assay.** The affinity and binding kinetics were measured using BIAcore assays. The dissociation rate constant (Kd) between His-ATF3 and JunB (Origene Technologies, Inc.) was determined using BIAcore 3000 (BIAcore, Inc.) and the data were analyzed using BIAEVALUATION software version 4.1 (BIAcore, Inc.). Standard ethyl(dimethylaminopropyl) carbodiimide/N-hydroxsuccinimide coupling was used to covalently immobilize JunB to CM5 sensor chips (BIAcore, Inc.) according to the manufacturer's instructions, and the interaction was detected in HBS-EP buffer (10 mM HEPES, pH 7.4, 0.15 M NaCl, 3 mM EDTA, 0.005% v/v surfactant P20). Flow cell 1 was left blank as a NC. Association rates were measured under a continuous flow of 10 µl/min using His-ATF3 at concentrations ranging between 30-150 nM plus a concentration of 0 nM, and the data were fitted using 1:1 Langmuir binding with no bulk refractive shift.

**In vitro His-tag pulldown assay.** Cell lysates containing His-ATF3 were prepared from the bacteria harboring pET32a-ATF3 plasmid after induction by IPTG for 4 h via sonication (0°C, 20 kHz, output 3; 50% duty cycle; 15 min) and incubated with 100 µl Ni-NTA agarose resin (Sigma-Aldrich; Merck KGaA) at 4°C overnight. After washing twice with PBS, bound proteins were boiled in the loading buffer, separated using 12% SDS-PAGE and detected by WB as aforementioned.

**Co-immunoprecipitation (Co-IP) assay.** Plasmids pCMV-Script-HA-ATF3 and pCMV-Script-Myc-JunB were prepared for the assay via inserting hemagglutinin (HA) or Myc sequence before the ORFs of ATF3 and JunB, respectively. For the Co-IP assay, HEC-1B-NC, HEC-1B-ATF3
or 293T cells (Chinese Academy of Sciences Committee Type Culture Collection) were transfected with 2.5 µg pCMV-Script-HA-ATF3 or pCMV-Script-Myc-JunB vector as aforementioned, respectively. After 24 h, cells were collected and then lysed in IP assay lysis buffer [10 mM Tris-HCl (pH 7.3), 100 mM NaCl, 1 mM EDTA, 0.2% Triton X-100, 0.2 mM dithiothreitol (DTT), 10% glycerol and protease inhibitors]. Cell lysates (1 mg) were incubated with 1 µg JunB antibody (1:50; cat. no. sc-8051; Santa Cruz Biotechnology, Inc.) or 1 µg immunoglobulin G1 (1:50; IgG1; cat. no. 401401; Biolegend, Inc.) as a control at 4°C overnight. The immunocomplex was precipitated with 25 µl Protein A Sepharose 6MB (Cytiva), pelleted at 800 x g, 4°C for 5 min, washed with 1 ml IP assay buffer supplemented with 300 mM NaCl three times and then detected using 12% SDS-PAGE and WB as aforementioned with anti-ATF3 antibody and anti-JunB antibody.

Co-localization assay. HEC-1B cells were seeded at a confluency of 80% on a glass slip and co-transfected with 5 µg pCMV-Script-HA-ATF3 and pCMV-Script-Myc-JunB plasmid, respectively, with Lipofectamine® 3000 (Invitrogen; Thermo Fisher Scientific, Inc.). The plasmids were prepared as aforementioned. After 24 h, cells were fixed in 4% formaldehyde for 15 min at 4°C, permeabilized with 0.2% Triton X-100 for 5 min at room temperature and blocked with 5% bovine serum albumin (cat. no. A7906; Sigma-Aldrich; Merck KGaA) for 50 min at room temperature. Cells were subsequently incubated with ATF3 (C-19) antibody (1:200; cat. no. sc-188; Sant Cruz Biotechnology, Inc.) and JunB antibody (1:200; cat. no. sc-8051; Santa Cruz Biotechnology, Inc.) at 4°C overnight and then labeled with FITC-conjugated anti-rabbit IgG (1:20,000; cat. no. 111-095-003; Jackson ImmunoResearch Laboratories, Inc.) and rhodamine (TRITC)-conjugated anti-mouse IgG (1:20,000; cat. no. 115-025-146; Jackson ImmunoResearch Laboratories, Inc.) for 1 h at room temperature. After washing with PBS three times, 30 sec each time, cells were counterstained with DAPI at room temperature for 10 min. After washing with PBS, cells were covered with Vectashield mounting media (Vector Laboratories, Inc.). Images were acquired with a laser scanning confocal microscope (Leica Microsystems, Inc.) using a x60 oil immersion lens.

Cell proliferation and colony formation assays. To assess cell proliferation, cells were seeded into 96-well plates at 2,000 cells per well. At the time points of 1, 2, 3, 4 and 5 days, the number of metabolically active cells was measured using a MTT assay (Sigma-Aldrich; Merck KGaA). With the formed formazan dissolved in 100 µl dimethyl sulfoxide, the absorbance values were measured at 490 nm with a Spectra Max 190 microplate reader (model 680; Bio-Rad Laboratories, Inc.).

For colony formation assays, cells were seeded in 6-well plates at 200 cells/well and grown in normal culture medium (DMEM/F12 with 10% FBS) at 37°C and 5% CO₂ for 9 days for colony formation. Colonies were fixed in 4% formaldehyde for 30 min at room temperature and then counted under a light microscope (magnification, x40; Eclipse Ts2; Nikon Corporation) after staining with 0.5% crystal violet for 20 min at room temperature. The experiments were repeated independently ≥3 times.

Chromatin IP (ChIP) assay. Cells with or without overexpression of ATF3 were fixed at room temperature in 0.5% formaldehyde for 20 min. After sonication (0°C; 20 KHz; output 2; 20% duty cycle; 5 min) using a sonifier (Model 450; Branson; Thermo Fisher Scientific, Inc.), IP was performed at 4°C overnight using 5 µg anti-ATF3 (1:50; cat. no. sc-87213; Santa Cruz Biotechnology, Inc.), anti-JunB antibody (1:50, cat. no. sc-8051; Santa Cruz Biotechnology, Inc.) or control mouse IgG1 (1:50; cat. no. 401401; Biolegend, Inc.). Pulled-down chromatin were denatured at 65°C overnight and the DNA was purified using the QIAquick purification column (size, 0.6 ml; Cytiva). The resulting AP-1 site included DNA was quantified by qPCR as aforementioned using the following primer pairs: Forward, 5'-TACTGTCTGGAAATCCCAC-3' and reverse, 5'-ACCCGGACTATCTGCCCAGGTC-3'. Fluorescence intensities were calculated using the following formula: Fluorescence intensity=(IP average ± IgG average)/(Input average).

Transwell migration and invasion assays. For Transwell migration assays, the lower surface of polycarbonate membranes of the inserts (8-µm pore size; Corning, Inc.) was coated with 100 µg fibronectin (Sigma-Aldrich; Merck KGaA). A total of 2x10⁴ cells suspended in 100 µl DMEM/F12 medium (without FBS) were dispensed into the upper chambers of the Transwells (with 0.6 ml DMEM/F12 medium supplemented with 10% FBS in the lower chamber) and incubated at 37°C for 24 h. Cells migrated to the lower membrane surface were fixed at room temperature for 20 min with 4% paraformaldehyde, stained with crystal violet (20 min at room temperature) and counted under a light microscope (magnification, x40; Eclipse Ts2; Nikon Corporation). For invasion assays, 10⁵ cells were plated in Transwells coated with Matrigel (1:6; BD Biosciences) at room temperature for 1 h and cultured at 37°C for 24 h. Invaded cells were stained and counted as aforementioned.

Apoptosis assay and cell cycle analysis. The apoptotic-related proteins were detected using WB as aforementioned. To measure the apoptosis ratios using flow cytometry, a total of 1x10⁵ HEC-1B-NC or -ATF3 cells were digested with 0.25% trypsin (without EDTA) and stained with FITC-Annexin V in fluorescence-assisted cell sorting buffer (cat. no. 556547; BD Biosciences) for 30 min at room temperature after washing with PBS. The total apoptotic (early and late apoptosis) cells were then analyzed with the software Kaluza version 1.3 (Beckman Coulter, Inc.) on a Beckman Gallios flow cytometer (Beckman Coulter).

For the cell cycle analysis, cells were fixed with 4% paraformaldehyde at room temperature for 2 h and stained with 7-aminoactinomycin D at room temperature for 10 min. After the cells were detected by flow cytometry (Beckman Gallios; Beckman Coulter, Inc.), cell cycle analysis was performed with the software Kaluza version 1.3 (Beckman Coulter, Inc.). Experiments were performed in as three independent replicates.

Caspase enzymatic activity measurement. The activities of Caspase-3/7, Caspase-8 and Caspase-9 were determined using the Caspase-Glo 3/7 Assay kit, Caspase-Glo 8 Assay kit and the Caspase-Glo 9 Assay kit (Promega Corporation),
respectively, according to the manufacturer's protocol. HEC-1B-NC, HEC-1B-ATF3, AN3CA-NC and AN3CA-ATF3 cells were seeded at a density of 4x10^5 cells/well in 6-well plates and cultured overnight. Following digestion, ~6x10^4 cells per sample were collected in 1.5-ml tubes. After 100 µl Caspase-Glo reagent (relative caspase substrate) was added to each sample, the cells were thoroughly mixed and incubated for 1 h in the dark at 37˚C. Luminescence values of each sample were measured using an IVIS Kinetic Imaging System (Caliper Life Sciences; PerkinElmer, Inc.). The quantitative analysis was performed based on the relative caspase activities and normalized to the raw luminescence units of the untreated control.

**JunB reporter assay.** JunB promoter constructs driving the expression of the luciferase reporter gene were used. A DNA fragment of the WT or mutant (MT) 5′ untranslated region (UTR) of JunB was cloned into the pRL-CMV luciferase reporter plasmid (50 ng; cat. no. E2261; Promega Corporation) and the resultant vectors were designated as ‘pRL-CMV-JunB-5′UTR-WT’ and ‘pRL-CMV-JunB-5′UTR-MT’, respectively. HEC-1B-NC or HEC-1B-ATF3 cells were transiently transfected with 1 µg Renilla constructs (as an internal control) or plasmids pRL-CMV-JunB-5′UTR-WT or pRL-CMV-JunB-5′UTR-MT using Lipofectamine® 3000 as aforementioned. Cell lysates were prepared after 24 h and the luciferase activities of cell lysates were then determined using a luminometer (Glomax 96; Promega Corporation). The relative luciferase activity was calculated as firefly/Renilla. All experiments were performed as ≥3 independent replicates.

**DNA precipitation (DNAP) assay.** Nuclear extracts (100 µg) in 500 µl nuclear extraction buffer (Thermo Fisher Scientific, Inc.) from HEC-1B-NC or HEC-1B-ATF3 cells were mixed with a biotinylated activator protein (AP)-1 site probe (50 pmol; cat. no. D3118; Beyotime Institute of Biotechnology) in a buffer [HEPES-KOH (pH 7.9), 80 mM KCl, 1 mM MgCl2, 0.2 mM EDTA, 0.5 mM DTT, 10% (w/v) glycerol, 0.1% Triton X-100 and 1 µg poly(dI-dC)] at 0˚C for 30 min. Then, 100 µl Streptavidin agarose beads (Sigma-Aldrich; Merck KGaA) were added and the mixture (0.6 ml) was gently agitated for 1 h at room temperature. Beads were then pelleted at 800 x g for 5 min at room temperature and washed three times with PBS, and the bound proteins were analyzed using WB and immunohistochemistry results (Fig. 1C).

**Xenograft assay in nude mice.** A total of 12 BALB/c nude mice (age, 4-5 weeks; weight 16-18 g) were purchased from Shanghai Laboratory Animal Research Center. The mice were housed in individually ventilated cages on a ventilated rack (Model GH; Suzhou Fengshi Laboratory Animal Equipment Co., Ltd.) in a specific pathogen-free facility with 12 h cycle of light/dark. The temperature, humidity and air flow of the ventilating air was set at 22˚C, 40% and 25 cm/sec, respectively. The animals were able to move freely with unlimited access to water and food. A total of 5x10^5 cells suspended in 100 µl 1X PBS were injected into the interscapular area subcutaneously of the mice. A group of mice (n=8) received HEC-1B-ATF3 (referred to as exATF3). Another NC group (n=4) received HEC-1B-NC cells transfected with control plasmid (cloning vector only, referred to as exNC). The sizes of tumors were measured daily over 4 weeks. Mice were sacrificed at 30 days post-injection by CO2 exposure at an air displacement rate of 30% per min. Tumors were excised and measured. The tumor volume (cm^3) was calculated by using the following formula: Volume=(longest diameter) x (shortest diameter)^2 x 0.5.

**Statistical analysis.** All statistical analyses were performed using SPSS 16.0 (SPSS, Inc.) or GraphPad Prism 5.0 (GraphPad Software, Inc.). Each experiment was performed ≥3 times and data are presented as the mean ± SD. A one-tailed, unpaired Student's t-test or Mann-Whitney U-test were utilized to determine significant differences between the treatment groups. P<0.05 was considered to indicate a statistically significant difference.

**Results**

**ATF3 is downregulated and JunB is upregulated in EC.** The expression levels of ATF3 and JunB were semi-quantitatively assessed using WB, indicating that ATF3 expression was low in EC tissues; it was >20-fold lower compared with healthy endometrium. However, JunB expression was relatively high, with a ~2-fold increase compared with healthy tissues (Fig. 1A). To further evaluate this result, healthy endometrial and EC tissues were subjected to immunohistochemical analysis. Strong staining for JunB and weak staining for ATF3 were observed in EC tissues (Fig. 1B). Thus, the results indicated that ATF3 was downregulated, while JunB was upregulated in EC.

Next, ATF3 and JunB expression levels were assessed in EC cell lines. WB analysis demonstrated that JunB was expressed in the type I and type II EC cell lines HEC-1B and AN3CA, while the expression of ATF3 was low, which was consistent with the WB and immunohistochemistry results (Fig. 1C).

**ATF3 overexpression impacts the proliferation, cell cycle, apoptosis, migration and invasion of tumor cells in vitro.** Following the identification of the low expression of ATF3 in ECs, HEC-1B cells were transfected to stably overexpress ATF3 in order to demonstrate the function of ATF3 in endometrial tumorigenesis. Stable transfection significantly increased the mRNA transcription of ATF3 by >180-fold (Fig. S1) and its protein expression by ~9-fold in HEC-1B cells (Fig. 2A). After transfection, the proliferation and colony-forming ability of HEC-1B cells was assessed using a MTT assay and a colony formation assay, respectively. The results suggested that overexpression of ATF3 in HEC-1B cells significantly inhibited cell proliferation (Fig. 2B) and colony formation (Fig. 2C) compared with exNC cells.

ATF3 is closely associated with the cell cycle, apoptosis, invasion and metastasis in certain tumor types (16-18). Therefore, it was investigated whether overexpression of ATF3 was able to change the cell cycle profile and trigger apoptosis in exATF3 cells. The flow cytometry results demonstrated that exATF3 cells had a markedly increased G1 phase population (60.9±3.0%; n=3), while the S (12.2±0.7%; n=3) and G2 phase (25.9±3.7%, n=3) populations were decreased, compared with those in exNC cells (41.2±3.2, 20.3±3.8 and 38.5±2.1%, respectively; n=3; Fig. 2D).
The cells of exNC and exATF3 were further analyzed with FITC‑Annexin V staining and were subjected to analysis of early and late apoptotic cell percentages using flow cytometry. The results indicated that a greater percentage of exATF3 cells (4.6±1.1%) were apoptotic compared with exNC cells (1.0±0.6%; Fig. 2E). In addition, cells transfected with ATF3 demonstrated a significantly reduced migratory and invasive abilities compared with the control cells (Fig. 2F). Collectively, the results suggested that ATF3 had a suppressive function in ECs.

Overexpression of ATF3 regulates the expression of apoptotic markers, MMPs and TIMPs. After the inhibitory effects of ATF3 on the proliferation, transmigration and invasion of ECs were identified, it was further investigated whether apoptotic signaling pathways were activated; thus, changes in the levels of apoptotic markers after overexpression of ATF3 in ECs were detected. The enzymatic activity of Caspases is regarded as an important indicator of apoptosis (19). Therefore, the activities of Caspase‑3/7, Caspase‑8 and Caspase‑9 were first measured, and it was found that the activities of all of these enzymes increased after ATF3 was overexpressed in the cells (Fig. 3A). In addition, the expression levels apoptotic-associated proteins, including cleaved PARP, Bax and p‑p38 increased, while Bcl2, and Bcl‑x decreased in the two cell lines (Fig. 3B). Thus, these results may explain the increased apoptotic ratios of host cells after overexpression of ATF3. Moreover, the relative protein levels of p‑p38 and p38 were found to be significantly increased, indicating that apoptosis may be mediated via a p38‑mitogen-activated protein kinase signaling pathway (Fig. 3B).

MMPs and TIMPs have prominent roles in cancer invasion and metastasis. TIMPs, binding to MMPs at a 1:1 molar ratio, are able to block the access of substrates to the catalytic domain of endopeptidases and eventually inhibit the bioactivities of MMPs (20). Furthermore, increases in the MMP/TIMP ratio have been reported to affect invasion and metastasis, and are widely considered as a marker for cancer invasion (21-23). Therefore, the mRNA expression levels of MMPs and TIMPs in both cell lines with and without ATF3 overexpression were compared. It was identified that certain MMPs, including MMP9, MMP10 and MMP13, were significantly downregulated following overexpression of ATF3 (Fig. 3C and D); MMP7 was slightly upregulated, but was not significantly different compared with the control. However, the expression levels of all TIMPs were not significantly different after ATF3 was overexpressed and remained stable.

Overexpression of ATF3 decreases the tumor size in a xenograft model. To further study the anti-tumor potential of ATF3,
a xenograft tumor growth assay in nude mice subcutaneously injected with stably transfected exATF3 or exNC cells was performed. The results indicated that overexpression of ATF3 significantly decreased the volume and weight of the xenograft tumors compared with the exNC group (Fig. 4A).

To determine the effect of ATF3 on proliferation and apoptosis, immunohistochemical staining of the tumor tissues for Ki67 and PCNA was performed. The expression of Ki67, indicative of tumor cell proliferation, was decreased by overexpressed ATF3 in the xenograft tumors. Moreover, notably weaker PCNA staining was detected in the exATF3 group compared with the exNC group (Fig. 4B).

Overexpressed ATF3 directly interacts with JunB and binds to AP-1 sites in ECs. The expression of ATF3 and JunB (Fig. S2) in EC was visualized by confocal microscopy. ATF3 and JunB were identified to be predominantly localized in the nucleus of EC cells (Fig. 5A). Previously, the binding
partners of ATF3 had been screened in a protein library of ECs using Y2H analysis and JunB was identified as one of its interactors (Fig. S3A and B). In addition, their interaction was demonstrated in vitro using Co-IP (Fig. S3C). To prepare the recombinant protein of ATF3, a prokaryotic expression plasmid, pET32a-ATF3, was constructed and His-ATF3 was expressed in bacteria. The bioactive His-ATF3 was obtained after affinity purification and denaturation. Using the His-ATF3, the interaction was assessed with a His-tag pulldown assay (Fig. S3D).

The presence of an ATF3/JunB interaction in ECs after ATF3 overexpression was assessed using Co-IP with monoclonal anti-ATF3 antibodies. In the native HEC-1B cells, no interaction between ATF3 and JunB was detected. However, from HEC-1B cells stably overexpressing ATF3, JunB was precipitated, indicating that the ATF3-JunB interaction only occurs in the ATF3-overexpressing cells (Fig. 5B).

To measure the affinity capacity between ATF3 and JunB, which represents the minimum concentrations of His-ATF3 or JunB required for binding, the association rate constant (Ka)
and Kd of the interaction between His-ATF3 and JunB were determined with a BLAcore instrument. Fitting the data to a 1:1 binding model yielded an apparent binding affinity with the equilibrium dissociation constant \( K_D = 1.9 \times 10^{-8} \) M, obtained from the ratio of the rate constants \( k_d / k_a \) (Fig. 5C).

Similar to the composition of the promoters of most MMPs (24,25), an AP-1 site is located in the promoter of JunB. WB analysis indicated that JunB was downregulated ~3-fold by overexpression of ATF3 (Figs. 5D and S4). To demonstrate that the downregulation of JunB was exerted by ATF3, luciferase plasmids with JunB promoter containing a WT AP-1 site or a MT AP-1 site at -2,322 to -2,315 were constructed (Fig. 5E). The highest luciferase activity was observed in the HEB-1C cells transfected with the plasmid containing the WT AP-1 binding site, and only slight or background bioactivities were measured when no AP-1 sites or a MT AP-1 site were included. In addition, the luciferase activity exhibited a significant decrease when ATF3 overexpression plasmid was co-transfected with reporter plasmid containing the WT AP-1 binding site. Furthermore, no difference in luciferase activity was identified between the groups transfected with exATF3 or exNC co-transfected with plasmid containing MT AP-1 binding sites or without promoter. Thus, the results indicated that the overexpression of ATF3 regulated JunB transcription and translation via binding to the AP-1 site in the promoter of JunB.

To demonstrate that the binding of ATF3 requires the presence of JunB, ChIP was performed. The JunB promoter DNA sequence was precipitated when either anti-ATF3 or anti-JunB antibodies were utilized for precipitation (Fig. 5F). In addition, the levels of precipitated DNA were in parallel with the expression levels of ATF3 or JunB. When DNA was precipitated with the AP-1 binding site, ATF3 and JunB were precipitated together along with the DNA, indicating that ATF3 regulated AP-1 signaling by interacting with JunB (Fig. 5G).

**Discussion**

EC is one of the most common cancer types of the female genital tract worldwide (1). While it is known that in the
majority of cases, estrogen and progesterone have a significant role in the pathogenesis of EC, the basic mechanism is yet to be fully elucidated. ATF3 has been reported to participate in the carcinogenesis of various types of cancer, such as prostatic...
the role of ATF3 in tumors is more complex and its influence on JunB in native ECs may be the low concentration of ATF3, which impedes the affinity between ATF3 and JunB. JunB complex. ATF3/JunB complex binds to the AP-1 site of the promoter of JunB to inhibit the expression of JunB, which mitigates cell proliferation, migration and invasion and facilitates cell apoptosis. X indicates transcriptional inhibition. ATF3, activating transcriptional factor 3; AP-1, activator protein 1; ORF, open reading frame.

Figure 6. Schematic diagram of the ATF3/JunB pathway in endometrial carcinoma, as a representative regulating model of AP-1 binding by to ATF3/ JunB complex. ATF3/JunB complex binds to the AP-1 site of the promoter of JunB to inhibit the expression of JunB, which mitigates cell proliferation, migration and invasion and facilitates cell apoptosis. X indicates transcriptional inhibition. ATF3, activating transcriptional factor 3; AP-1, activator protein 1; ORF, open reading frame.

cancer (26), but the roles of ATF3 in EC remain unknown. The present results demonstrated that ATF3 inhibited the proliferation, migration and invasion of EC cells, at least in part by interacting with JunB.

JunB is an important transcriptional factor regulating the cell cycle, thus reflecting cell proliferation in ECs (27,28). JunB has been shown to have an oncogenic role and can promote cancer progression, invasion and metastasis in uveal carcinoma (29), multiple myeloma (30), breast cancer (31) and squamous cell carcinoma (32,33). Furthermore, JunB acts as an oncogene and induces abnormal proliferation of previously quiescent cells (34). JunB, as a member of the AP-1 family, is considered to exert its function via AP-1 signaling (35). Various mechanisms underlying the AP-1-mediated regulation of tumor progression, particularly invasion and metastasis, have been reported. For instance, JunB regulates several genes, including MMP2, MMP9 and C-C motif chemokine ligand-2, to enhance tumor invasion and angiogenesis in renal cell carcinomas (36). With AP-1 sites in the promoters, MMPs are considered to be regulated by AP-1, formed by AP-1 family members, including JunB (24,25). Therefore, JunB may be a promising therapeutic target for various cancer types.

In the present study, it was demonstrated that ATF3 was able to impair the function of AP-1 and JunB to regulate MMP by forming a complex with JunB. In addition, ATF3 suppressed the transcription and translation of JunB (Fig. 6). To the best of our knowledge, this is a novel mechanism regarding the regulation of the function of JunB in carcinomas. In addition, the minimum concentration requirements for the interaction between ATF3 and JunB were examined, and it was speculated that the possible reason as to why ATF3 did not bind to JunB in native ECs may be the low concentration of ATF3, which impedes the affinity between ATF3 and JunB.

Similar to JunB, ATF3 is a member of the AP-1 family, but the role of ATF3 in tumors is more complex and its influence on cells is context-dependent (37). ATF3 regulates gene expression in various conditions via binding to the different ATF3/cyclic adenosine monophosphate-responsive element-binding protein cis-regulatory elements (38-40). Moreover, the expression of ATF3 is activated by transforming growth factor-β binding to its receptor, resulting in the phosphorylation of SMAD3. The stress signal, acting via p38 kinase, is also able to induce ATF3 expression (41).

ATF3 can inhibit apoptosis induced by stimuli in cardiac myocytes (42), but it also induces apoptosis in tumors (16). Moreover, in varying tumor types, the functions of ATF3 may be different or even the opposite. Under certain circumstances, ATF3 has been shown to promote oncogenesis and migration in skin, breast, prostate and colon cancer (17,43-46). ATF3 expression may be associated with increased metastasis in melanoma and in breast cancer cells (47,48), and knockdown of ATF3 reduces the ability of HT29 colon cancer cells to invade (46), suggesting that ATF3 acts as a cancer promoter. However, previous studies on ATF3 have also revealed a tumor suppressor role in colon, breast, prostate, bladder, glioblastoma and lung cancer by inhibiting cell proliferation and metastasis, and promoting cell death and apoptosis (47-53). Furthermore, a previous study reported that ATF3 may function as a tumor suppressor in Ras-mediated tumorigenesis (54). It has been suggested that the various and context-dependent roles of ATF3 in cancer are due to the complex ATF3-associated protein-protein interaction networks. Thus, ATF3 may be capable of interacting with numerous important and critical proteins to regulate their functions, in addition to its transcriptional regulation (55).

The ATF3/JunB interaction was previously identified to exert an inhibitory effect on human prostate cancer (45). In line with this finding, the present results demonstrated that ATF3 was able to inhibit the progression, invasion and metastasis of ECs via binding to the AP-1 sites in the promoters of various genes, including JunB and MMPs. However, it remains elusive whether ATF3, independent of the ATF3/JunB complex, is able to help degrade MMPs via a proteasome-dependent pathway, as has been reported in esophageal cancer (56). The present findings may explain why ECs lost their transmigration and invasion ability after ATF3 overexpression, as the ratio of MMPs/TIMPs, regulated by ATF3, decrease significantly. Moreover, these results are in line with a previous study, which revealed that ATF3 was able to reduce the ability of human glioblastoma to migrate by regulating MMPs and TIMPs (21).

In our previous study, Y2H mating experiments of ATF3 were performed and >274 interactions were identified in human healthy ovary tissues, of which JunB is one of the most positive interacting partners of ATF3 (data not yet published). This finding is consistent with the result of the present study and the earlier studies, which demonstrated that the Fos/Jun and ATF/cAMP response element binding protein families of transcription factors function in dimerization (57), which is mediated by the ‘leucine-zipper’ motif, and bind to DNA to alter the expression of specific target genes. While the interaction between ATF3 and JunB has been revealed in other tissues or cell types, to the best of our knowledge, the present study was the first to assess this interaction in ECs. The results suggested that the inhibition of ECs by ATF3 may be used as a novel therapeutic strategy for EC.
In conclusion, analysis of EC tissue samples from 50 cases identified a weak expression of ATF3, but upregulation of JunB expression in EC tissues. It was found that overexpression of ATF3 resulted in downregulation of JunB expression. Further investigation demonstrated that the ATF3/JunB complex bound to the promoter and inhibited the expression of JunB, thus resulting in the remodeling of EC cells and mitigated developments of EC in vitro and in vivo. Therefore, JunB may serve as a potential therapeutic target for EC, and ATF3 as an effective therapeutic tool.

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Availability of data and materials

The datasets used or analyzed during the current study are available from the corresponding author upon reasonable request.

Authors’ contributions

FW, JG, HW and JL conducted the experiments and analyzed the results. FW and JG conceived the experiments. FZ and JG supervised the project and interpreted the data. FW, FZ and JG wrote the article. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The protocol (approval no. 0101N16032) of the present study was approved by the Human Investigation Ethical Committee of Shanghai First People’s Hospital Affiliated Shanghai Jiao Tong University (Shanghai, China) and Renji Hospital Affiliated to Shanghai Jiao Tong University School of Medicine (Shanghai, China). All samples were obtained from patients who had provided written informed consent for the use of their tissues for the purposes of research after the operation.

The animal experiment was in accordance with the recommendations in the Guidelines for the Care and Use of Laboratory Animals of China (http://www.nsfc.gov.cn/nsfc/cen/pfzl/pufanew/20110801_02.pdf). The protocol (approval no. 2013020346) was approved by the Committee on the Ethics of Animal Experiments of the School of Pharmacy of Shanghai Jiao Tong University (Shanghai, China).

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

References


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